The State of Standards and Interoperability for mHealth among Low- and Middle-Income Countries

mHealth Alliance

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The State of Standards and Interoperability for mHealth among Low- and Middle-Income Countries
Executive Summary

The mHealth Alliance commissioned this report to review the state of health informatics standards and interoperability with respect to the use of mobile health among low- and middle-income countries (LMICs) with the aim of identifying critical gaps and opportunities to support the scale up of mHealth. The intended audience for this report is stakeholders developing or employing health informatics standards, and implementers and vendors seeking to interoperate systems. This report provides guidance for how the above stakeholders may align efforts around the use of standards to achieve interoperable mHealth at scale.

This report is based on three sources of information: (1) a landscape scan of current initiatives and gray literature related to mHealth standards and interoperability; (2) a review of academic articles examining the use of standards to achieve interoperability within mHealth; and (3) unstructured informal interviews with 33 informants who directly or indirectly influence the uptake of health informatics standards and the structure of the national ICT environment.

Improved continuity of care across provider, place, and time is one of the great promises of mHealth, but this can only be achieved if information is readily available at the right place and time, regardless of how or on what information system it was collected. Interoperability of data in such heterogeneous environments is enabled through the application health informatics standards within the context of a strategic eHealth framework. Such a framework must address critical issues of governance, infrastructure, architecture, workforce capacity, policy, and financing.

There are many organizations actively developing standards and standards implementation guidelines to achieve interoperability within a specific health domain, such as the International Organization for Standardization (ISO), Health Level 7 International, and Integrating the Healthcare Enterprise. The World Health Organization and International Telecommunications Union play both normative and informational roles with respect to standards development and dissemination. While standards development organizations (SDOs) have historically focused on high-income country health systems, due to greater readiness to adopt and substantial government and private industry investment, there is a notable broadening of scope to accommodate LMIC needs, as evidenced through the formation of LMIC-focused working groups and the 2012 ISO publication of the Capacity-based eHealth Architecture Roadmap.

Yet significant gaps exist in the capacity and participation of LMICs in the development and adoption of health informatics standards. The largest challenges are lack of local informatics human resource capacity and insufficient investment into standards activities, which prevents productive engagement with international standards development organizations (SDOs).

mHealth is a subset of eHealth and, as such, many of the issues surrounding the scale up and harmonization of eHealth are the same for mHealth, such as the need for national eHealth strategies and stronger eHealth governance. mHealth within LMICs also presents unique challenges, including limited bandwidth and infrastructure, reliance on insecure or unreliable messaging protocols such as the short messaging service (SMS), and the shift towards non-clinical and community-based workers delivering new types of services and in new settings.
While the body of academic literature related to mHealth and standards has grown over the last decade, the literature review indicated that there have been no evaluations of the comparative advantages of using health informatics standards to achieve interoperability of eHealth and mHealth systems focused on LMICs. However, at the time of publication, there were very few scaled-up and interoperable implementations to be evaluated, so this result was not unexpected. LMIC-focused articles were generally descriptive in nature, often advocating for global alignment around standards, interoperability, and capacity issues. Studies assessing the comparative advantages of specific video compression and security protocols have been published and should be applied to future telemedicine deployments. There remains a gap in the evaluation of the use of specific packages of standards to achieve specific health system or health information system goals.

Key informants noted a recent shift in the global eHealth discourse, in that most LMICs have developed or are developing national eHealth strategies and that interoperability was nearly universally accepted as a key element of these strategies. Governments were considered to have the most important role in influencing the design and requirements of mobile and electronic health systems implemented in each of their countries, and local implementers and development partners generally wanted a more robust and enabling regulatory environment in place to provide this guidance and lower investment risk. Donors were viewed as playing a pivotal role in providing financial incentives to shift the market in favor of interoperable systems. Key informants acknowledged that health informatics workforce capacity was lacking among all stakeholders aside from SDOs, but that SDOs generally had little experience in LMICs.

The gaps and opportunities with respect to the use of standards to achieve interoperable mHealth at scale can be categorized into five buckets: (1) Building capacity and increasing standards access; (2) Filling in standards gaps for adoption among LMICs; (3) LMIC engagement and mediation; (4) Promoting coordination and alignment; and (5) Strengthening mHealth and eHealth governance. More details about each of these categories are provided below.

1. All stakeholders other than SDOs generally lacked capacity in health informatics expertise, although the shortage was most severe among LMIC governments and local implementers. Capacity deficiencies may be addressed by establishing formal health informatics training and education programs, better informing local decision and policy makers of the strategic implications of adopting standards and interoperability best practices, improving donors and implementers’ ability to strategically invest in an interoperable health system, empowering local professionals and professional organizations, and addressing barriers to standards access.

2. Most existing health informatics standards can be repurposed without alteration in LMIC contexts, although adaptation and possibly new standards development may be necessary in some cases in order to support unique LMIC business requirements and new mHealth use cases. The development of a mHealth reference architecture will provide critical guidance on the design of mobile health information systems that is consistent and compatible with other reference architectures. Implementation guides are needed to assist LMIC governments and implementers in the adaptation of standards for local contexts.
3. Bridging the standardization gap in the short-term means ensuring that LMIC requirements are represented in health informatics standards. Implementers and development partners with experience working in specific regions can and should, in partnership with governments and SDOs, collate and contribute LMIC requirements to the standards development process to ensure that the final product is applicable globally. The long-term solution to the standards gap is for LMICs to engage productively in the standards development process as participating members. Regional eHealth standardization collaboratives, established by local governments and in partnership with academic institutions, private sector, SDOs and other international organizations, may be an effective mechanism for joint regional learning, engagement with the international standards community, and a valuable step on the way towards forming independent national standards bodies.

4. Mechanisms to promote coordination and alignment between governments and their partners are needed to broker the social and institutional relationships necessary for interoperability among and between other stakeholder types. Guidelines should be developed to help donors strategically invest in technology implementations and platforms that align with country priorities and international best practices. Regional eHealth standardization collaboratives could serve as an effective forum for identification and development of local priorities and standards requirements. Methods should be explored for requiring standards compliance in grants, possibly through certification, which would serve to shift market incentives to favor interoperability.

5. As mobile health systems move towards scale, existing guidelines and strategies will need to be revised to reflect new demands on (1) Executive sponsorship; (2) National leadership of eHealth programs; (3) eHealth standards adoption and implementation; (4) Development of eHealth capability and capacity; (5) eHealth financing and performance management; and (6) eHealth planning and architecture maintenance. Regulations regarding standards adoption and interoperability of mHealth and eHealth systems will play a key role in shifting market dynamics to favor interoperability.

Three crosscutting recommendations were identified that are intended to frame subsequent dialogue around near-term opportunities for the global community to address these challenges in a more coordinated fashion:

1) Shift market dynamics to incentivize interoperability, primarily through regulatory and financing mechanisms;

2) Align investments and implementations with national and regional eHealth strategies; and

3) Establish regional eHealth standardization collaboratives to facilitate pooling of resources and expertise and to serve as focal point for standards activities in a region.
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<th>Acronym</th>
<th>Description</th>
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<tr>
<td>CEN</td>
<td>European Committee for Standardization</td>
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<tr>
<td>CHW</td>
<td>Community Health Worker</td>
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<td>DICOM</td>
<td>Digital Imaging and Communications in Medicine</td>
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<td>EHR</td>
<td>Electronic Health Record</td>
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<td>ESA</td>
<td>European Space Agency</td>
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<td>FLW</td>
<td>Front Line Worker</td>
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<td>GOe</td>
<td>Global Observatory for eHealth</td>
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<td>GPRS</td>
<td>General Packet Radio Services</td>
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<tr>
<td>GPS</td>
<td>Global Positioning System</td>
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<tr>
<td>GSMA</td>
<td>Groupe Speciale Mobile Association</td>
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<td>HIS</td>
<td>Health Information System</td>
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<td>ICT</td>
<td>Information and Communications Technology</td>
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<td>IEEE</td>
<td>Institute of Electrical and Electronics Engineers</td>
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<td>IHE</td>
<td>Integrating the Healthcare Enterprise</td>
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<td>ISO</td>
<td>International Organization for Standardization</td>
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<td>ITU</td>
<td>International Telecommunications Union</td>
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<tr>
<td>LMIC</td>
<td>Low- and Middle-Income Country</td>
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<tr>
<td>MNO</td>
<td>Mobile Network Operator</td>
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<td>NGO</td>
<td>Non-Governmental Organization</td>
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<td>NIH</td>
<td>US National Institutes of Health</td>
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<td>SDO</td>
<td>Standards Development Organization</td>
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<td>SMS</td>
<td>Short Messaging Service</td>
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<td>UN</td>
<td>United Nations</td>
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<td>USAID</td>
<td>United States Agency for International Development</td>
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<td>USSD</td>
<td>Unstructured Supplementary Service Data</td>
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<td>WHO</td>
<td>World Health Organization</td>
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Background

Mobile health, or “mHealth,” refers to the use of mobile telecommunication and multimedia technologies to support service delivery and health systems. mHealth is a subset of eHealth, which the World Health Organization (WHO) defines as the use of information and communications technologies for health. Mobile technology is being used increasingly to augment health service delivery throughout the world. As mHealth projects scale up, integration of mobile health programs and applications into the broader health information system becomes essential.

Even in high-income economies, health systems are struggling to realize the full potential of information and communications technology (ICT) to strengthen health systems and support care delivery, due, in part, to limited technological and institutional interoperability. The promise of a connected and interoperable health system is that regardless of how data is collected, the right information is available to the right person, in the right place, and at the right time. With the advent of mobile health, there is an even greater need to ensure reliability of services, availability of information, and patient privacy and confidentiality. In particular, continuity of care across provider, time, and institution requires that information is shared, trusted, and secured. Low- and middle-income countries (LMICs) have perhaps the greatest potential to extend health systems’ reach by using mHealth to integrate remote workers and patients into the health system. Interoperability in such heterogeneous environments necessitates improved governance and alignment at the national and international levels, and greater diffusion of existing health informatics standards, adaptation of standards to LMIC contexts, and, in some cases, development of new standards.

Standardization at the national level can only be achieved by aligning efforts of international and bilateral institutions, standards development organizations (SDOs), donors, mobile network operators (MNOs), and other private sector actors with local government and health system priorities. The standardization gap means that representatives from developing countries are not at the table contributing their unique requirements for the use of ICT or standards utilization to the standards development process. Furthermore, even if LMIC requirements are represented in health informatics standards, there is a large gap between development of a standard and effective adoption of that standard. High cost of standards adoption, lack of prioritization, and limited local health informatics capacity remains a significant challenge in many countries.

While mHealth-enabled interventions have been documented in countries with eHealth ecosystems at all levels of maturity, interventions expected to have the greatest impact rely on more advanced ICT infrastructure. For example, countries with limited ICT infrastructure and expertise may be able to leverage mobile networks’ short messaging service (SMS), or text messages, to send automated alerts and appointment reminders to remote health workers. But, the same countries may find that network reliability prohibits expansion of telemedicine services beyond city centers.

The WHO and International Telecommunications Union (ITU) describe the national context for eHealth development in terms of the ICT environment, which represents the physical infrastructure and market adoption of components that utilize the infrastructure, and the ICT enabling environment, which consists of governance, policy, standards, human resources, and other aspects that are critical in the scaling of ICTs (Figure 1).
National Context for eHealth Development

Figure 1. WHO-ITU National eHealth Strategy Toolkit
Numerous global health and health informatics organizations have cited interoperability as a major obstacle to the meaningful use of ICTs at scale (see references 2, 5, 27). Development of a national standards and interoperability framework provides the context and forum for the multi-sectoral engagement necessary to address these issues in a deeply interconnected health system. Guidelines for this process are described in several key documents, including the 2012 National eHealth Strategy Toolkit developed by the WHO and the ITU.4

The mHealth Alliance Business Plan5, developed with Dalberg in 2011, identified limited technological integration and interoperability amongst mobile and electronic health systems as a key barrier to scaling mHealth. The report further identified key issues that contributed to limited technological integration and interoperability:

1) Lack of alignment on standards
2) Limited opportunities for collaboration
3) Limited collaboration or communication across other mServices (e.g. mobile banking, mobile education)
4) Limited guidance from donors and governments on which standards to use

The mHealth Alliance commissioned this follow-up report to identify critical gaps, as well as opportunities, with respect to the current state of health informatics standards and interoperability to support the scale of mHealth systems. The audience for this report includes stakeholders developing or employing health informatics standards, policy makers creating eHealth guidelines, national standards bodies or professional organizations, and implementers or vendors seeking to interoperate systems.

Findings are broken into the following sections:

1) a conceptual framework for approaching the complex space of standards, technical integration, and interoperability;
2) an assessment and catalogue of key obstacles that are preventing mHealth from achieving scale;
3) a review of published and gray literature related to the use of health informatics standards and interoperability within mHealth in LMICs;
4) results and analysis of key informant interviews; and
5) a discussion of opportunities and strategies for addressing key obstacles.
Methodology

Information contained in this report was derived from three sources:

1) Landscape scan of current initiatives and gray literature produced by multi- and bi-lateral institutions, governments, and major implementing organizations, etc;

2) Published literature review of peer-reviewed journals; and

3) Key informant interviews of individuals representing a cross-section of institutions and stakeholders involved in the scale up of mobile health programs, including donors, policy makers, mobile network operators, academics, implementers, and technology developers.

While every effort was made to present a balanced and objective view of the current state of standards and interoperability with respect to mHealth, there is a bias and subjectivity that is both unavoidable and useful for representing stakeholders’ diverse engagement with the field.

Literature Review

The literature review consisted of a search in PubMed for scientific published articles that contained both a “mobile health” term and a “standards” or “interoperability” term. References in the articles were followed to identify additional relevant materials. “Mobile health” search terms included:

- mHealth
- mHealth
- mobile health
- mobile phones
- mobile technology
- telemedicine
- telehealth

The "standards" and "interoperability" search terms included:

- standardization
- standard(s)
- interoperability
- integration
- architecture

No crosscutting systematic reviews of mHealth standards and interoperability were found, which would have provided a lower time bound for the literature search. Therefore, the search was restricted to articles published in the year 2002 or later based on the assumption that this timeframe would capture the vast majority of articles in the still nascent field of mobile health.
A total of 2,154 published articles from peer-reviewed sources were identified in PubMed and by following references. All articles were then screened for their relevance to the subject of this report. The exclusion criteria used for screening were as follows:

- Literature proposing a reference architecture without specific mention of mobile health were excluded. Articles describing specific clinical interventions that used mobile devices, evaluations of mobile health interventions measuring health outcomes, or particular software products or platforms were excluded, unless addressing particular standards or interoperability issues. Articles covering the “quantified self,” implanted or embedded sensors and sensor networks, implantable devices, social or peer-to-peer networks, grid networks, or public health databases such as genetics or surveillance were also excluded. Articles discussing standards related to the health effects of electromagnetic radiation from mobile devices were also excluded.

- Articles that use “telemedicine” to refer to the use of internet-based video conferencing to connect a clinician to a remote patient were excluded, unless standards were a primary focus, as were literature and programs that leveraged information and communication technology infrastructure to link traditional computers as opposed to mobile devices.

Note that while this literature review was methodical, it was not a systematic review in the strict sense. Also note that full versions of articles were not accessed unless available for free or judged to contain relevant insights not available in the abstract alone.

**Key Informant Interviews**

A total of 33 key informants were interviewed by phone or in person between May 8 and September 3, 2012. Informants represent a non-random convenience sampling of a broad range of stakeholders engaged in mHealth standards and interoperability. Interviewees were categorized as follows:

- Mobile Network Operators [1]
- Standards Development Organizations [4]
- NGOs / Implementers [4]
- Technology Developers [9]
- Researchers / Academics [3]
- Donors [5]
- Multinational Organizations [5]
- Government [2]

Of the key informants, 5 [15%] were female, and 28 [85%] were male. 15 [45%] were US-based, and 7 [21%] represented LMICs. Refer to Appendix A for the full list of interviewees.
Overview of Health Informatics Standards and Interoperability

The field of health informatics standards and interoperability is expansive and touches on hardware, infrastructure, software, semantics, social and institutional structures, and the interactions between each of these. This section describes the stakeholders involved in the development and harmonization of health informatics standards, provides working definitions for standards and interoperability, and presents a comprehensive eHealth architecture as proposed by the International Organization for Standardization (ISO).

Stakeholder Map

There are a number of organizations developing or contributing to the development and use of health informatics standards globally. Several key organizations are listed below:

- **ISO Technical Committee 215 (ISO TC 215)** develops standards across all areas of health informatics. It is divided into working groups that focus specifically on data structures, architecture, interoperability, devices, privacy and security. TC 215 is the governing body of the Public Health Task Force (PHTF), which was established to improve the uptake of health informatics standards among low-income countries.6

- **Health Level 7 International (HL7)** develops standards for the interoperability of health information technology. Examples include the HL7 Electronic Health Record Functional Model and the HL7 Clinical Document Architecture.

- **European Committee for Normalization (CEN) TC 251** is the health information technology technical committee of the European Union SDO. ISO TC 215 and CEN TC 251 regularly work jointly in standards development efforts.

- **CDISC (Clinical Data Interchange Standards Consortium)** develops standards for the interoperability of medical research information and related systems. Examples of standards include the Study Data Tabulation Model (SDTM), Standard for Exchange of Non-clinical Data (SEND), and Analysis Data Model (ADaM).

- **GS-1** develops standards related to supply-chains and their management. The GS-1 System is comprised of standards for Barcodes for identification, eCom for business messaging standards, GDSN (global data synchronization network), and EPCGlobal for tracking.

- **IHTSDO (International Health Terminology Standardization Organization)** owns and maintains SNOMED CT, which stands for Systematized Nomenclature of Medicine – Clinical Terms. This is the most comprehensive terminology dictionary for diseases, clinical findings and symptoms, procedures, etc.
• **IHE (Integrating the Healthcare Enterprise)** is an industry and professional organization working to improve information sharing between healthcare information systems. Note that IHE is not a SDO, rather it creates profiles for developers and users, such as the Cardiology or Laboratory profiles, to guide implementers and vendors on standards compliance necessary to achieve a specified level of interoperability within a domain.

The seven organizations above comprise the Joint Initiative on SDO Global Health Informatics Standardization, commonly known as the **Joint Initiative Council (JIC)**. The JIC was organized to facilitate coordination and harmonization of standards development activities. The JIC provides an important mechanism for joint publication of standards by two or more SDOs and a forum for resolving conflicts.

There are several organizations that play important roles in the field of health informatics that are not directly linked to the JIC:

• **International Telecommunications Union (ITU)** is the United Nations agency responsible for information and communications technology. ITU has specific arms for Standardization (ITU-T) and for the Development Sector (ITU-D).

• **World Health Organization (WHO)** plays an important leadership role in both standards development and in interoperability priority setting and standards dissemination. The WHO maintains and produces the WHO Family of International Classifications (FIC) to provide a common, global framework and language for health information. The FIC includes the International Classification of Diseases (ICD), International Classification of Function, Disability and Health (ICF), and International Classification of Health Interventions (ICHI). The WHO also developed the standard for Statistical Data and Metadata Exchange – Health Domain (SDMX-HD) for exchange of monitoring and evaluation data. The WHO also has several entities that interact with the standards space through leadership and prioritization, guideline development, and convening. These include the Department of Knowledge Management and Sharing, Department of Health Statistics and Health Information, and the Global Observatory for eHealth (GOe), which provides valuable information to support governments on the strategic adoption of electronic health information systems. GOe has published reports on key eHealth topics, including eHealth country profiles, telemedicine, mHealth, safety and security, legal frameworks, and management of patient information.

• **International Medical Informatics Association (IMIA)** is a member organization for national or regional informatics associations that promotes the effective use of informatics within healthcare, the dissemination of knowledge and health informatics education, and translation of research into practice. IMIA has significant representation throughout the world, including LMICs. IMIA Working Groups on Standards, Health Informatics for Development and Education deal directly with eHealth in LMIC countries.

• **Public Health Data Standards Consortium (PHDSC)** is a US-based member organization that aims to empower healthcare and public health communities with health information technology standards to improve individual and community health. PHDSC contributes to international standards development efforts and is engaged in development of standards curricula.
- **Continua Health Alliance** is a non-profit industry organization of more than 220 healthcare and technology companies worldwide focusing on interoperable personal connected health devices and solutions. Continua provides product certification and industry recommendations to SDOs.

- **African Union eHealth Expert Group** is a member project that is making recommendations for the harmonization of eHealth across the African continent and drafting the document “African Union eHealth Development to Support the Africa Health Strategy: 2007-2015”. This group represents many of the key local health informatics experts.

In 2005, the WHO General Assembly adopted a resolution promoting the use of eHealth to achieve health system goals among all member states. The Regional WHO Committee for Africa adopted a similar resolution in 2010, with key recommendations for African states:

- Establish national eHealth strategies and implement monitoring and evaluation systems to track progress
- Strengthen national political commitment, leadership and coordination by establishing intersectoral support mechanisms for eHealth
- Establish a conducive policy environment
- Build out infrastructure and establish services for eHealth (e.g. Internet connectivity and online resources)
- Develop human capacity for eHealth by establishing sustainable education and training opportunities
- Set up sustainable financing for eHealth infrastructure and ICT integration, and encourage private sector engagement

Several global initiatives build off the above commitments across multiple programmatic areas. The Commission on Information and Accountability for Women’s and Children’s Health was established in 2011 to “improve global reporting, oversight and accountability for women’s and children’s health.” Integration of ICTs into national health information systems and adoption of technical standards are key recommendations of the commission. The Innovation Working Group and *Every Woman Every Child* movement are investing into mobile and electronic tools to accelerate progress towards reductions in maternal and child mortality. Innovative technologies, including mobile phones and SMS, are also a key focus of the UN Commission on Life-Saving Commodities for Women and Children’s Health for their role in facilitating access to selected products and in complementing and informing service delivery. The WHO is garnering support for a global resolution specifically on eHealth standards and interoperability with the goal of adoption and harmonization.

The health informatics standards community has begun to mobilize resources to increase the accessibility of standards, to address limited adoption of standards among LMICs, and to fill mHealth standards gaps. The Standards Knowledge Management Tool (SKMT Glossary), supported by the Joint Initiative Council, is a free web-based portal that serves as a single point of reference for all published standards and definitions from JIC members and other relevant documents such as WHO publications. The Health Informatics Wiki combines information from national eHealth case studies with the SKMT Glossary to

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1 [www.skmtglossary.org](http://www.skmtglossary.org)
assist in identification of standards relevant to a particular health system context. Several efforts, including the Public Health Task Force at ISO and the Africa Initiative of HL7’s International Mentoring Committee, specifically aim to increase standards capacity and adoption among LMICs. HL7, IHE, and HIMSS have formed distinct mHealth task forces or working groups to address unique standards gaps related to the use of mobile and smart devices within health care. SDO efforts are expected to result in the development of a reference architecture and domain profiles for using mHealth in health systems at various levels of maturity and health domains, and integrating it with other health information systems, including electronic health records, laboratory information management systems, or district health information systems.

**Health Informatics Standards**

ISO defines a standard as the following:

“A standard is a document that provides requirements, specifications, guidelines or characteristics that can be used consistently to ensure that materials, products, processes and services are fit for their purpose.”

There are several ways to classify and scope health informatics standards. For the purposes of this report, we use the divisions of ISO Technical Committee 215 on Health Informatics (ISO TC 215), which are broken into working groups as follows:

- **Architecture (Working Groups 1 and 8)** – Definitions, templates, data sets. Standardized models of the functions and conformance criteria associated with a system that is platform agnostic (e.g., HL7 Electronic Health Record System Functional Model), including definition of business requirements and data structures.
- **Data interchange (Working Groups 2 and 7)** – Harmonization and adaptation of clinical and administrative messages for system and device interoperability.
- **Semantic content (Working Group 3)** – Develops standards for the representation of terminological resources, such as health concepts and data.
- **Security, safety and privacy (Working Group 4)** – Confidentiality, integrity and availability, accountability, security management, and information systems safety.
- **Pharmacy and medicines business (Working Group 5)** – Develops standards for interoperability of e-pharmacy systems and other medicines applications.

It is important to clarify that ISO standards are considered “open” according to the following definition: a) the entire contents of each standard may be obtained; b) each standard was developed and defined as part of an open, transparent process; and c) ISO standards are not subject to sudden changes as modifications are sent through a pre-defined systematic review. However, “open” ISO standards are generally not free.

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2 www.hiwiki.org
Figure 2. ISO, HL7 and IMIA membership categorized by country economic grouping as of September 2011.

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<thead>
<tr>
<th>ECONOMIC GROUPING</th>
<th>ISO TC 215 P</th>
<th>ISO TC 215 O</th>
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<td>-</td>
<td>Burundi, Malawi, Mali, Togo</td>
</tr>
<tr>
<td>LMIC</td>
<td>Armenia, Honduras, Philippines</td>
<td>Mongolia, Ukraine</td>
<td>-</td>
<td>Cameroon, Côte d'Ivoire, Ghana, Honduras, Nigeria, Pakistan, Philippines, Sri Lanka, Ukraine</td>
</tr>
<tr>
<td>UMIC</td>
<td>Brazil, China, Iran, Malaysia, Mexico, Russian Federation, South Africa, Tunisia, Turkey</td>
<td>Argentina, Bulgaria, Colombia, Ecuador, Kazakhstan, Germany, Romania, Serbia, Thailand</td>
<td>Argentina, Bosnia and Herzegovina, Brazil, China, Colombia, Mexico, Romania, Russian Federation, Turkey, Uruguay</td>
<td>Argentina, Bosnia and Herzegovina, Brazil, Chile, China, Cuba, Iran, Mexico, Peru, Romania, South Africa, Thailand, Turkey, Uruguay, Venezuela</td>
</tr>
</tbody>
</table>

Figure 3. LIC, LMIC, UMIC standards organization membership.

<table>
<thead>
<tr>
<th>GEOGRAPHIC REGION</th>
<th>COUNTRIES</th>
<th>ISO TC 215 P</th>
<th>ISO TC 215 O</th>
<th>HL7</th>
<th>IMIA</th>
</tr>
</thead>
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<tr>
<td>Sub–Saharan Africa</td>
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<td>1</td>
<td>0</td>
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<tr>
<td>South Asia</td>
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<td>0</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Europe &amp; Central Asia</td>
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<td>5</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>East Asia &amp; Pacific</td>
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<td>2</td>
<td>1</td>
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<tr>
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<td>8</td>
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<td>Middle East &amp; North Africa</td>
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<td>2</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Other</td>
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<td>11</td>
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<td>57</td>
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</tbody>
</table>

Figure 4. ISO, HL7 and IMIA membership categorized by geographic region.
The Standardization Gap

Significant gaps exist in the capacity and participation of LMICs in the development and adoption of health informatics standards. This is due to an interrelated set of factors, including: lack of local informatics human resource capacity and high complexity of standards; low prioritization by governments, vendors, and implementers; limited access to existing standards; limited funding for engagement with international SDOs; absence of national standards bodies; and inadequate ICT infrastructure for standards participation. These challenges, as well as potential solutions, are laid out in the 2009 ITU-T Bridging the Standardization Gap.3

Low- and middle-income countries have limited representation and membership in international standards organizations, which limits stakeholders’ ability to contribute their unique requirements to the standards development process. Within ISO TC 215, there are two membership types: Participating members (P-members) are full members and are requested to vote on every proposed standard; Observing members (O-members) may contribute but are not permitted to vote. Figure 2 illustrates the marked difference in representation of low-, lower-middle-, upper-middle-, and high-income country membership in ISO TC 215, HL7 International, and IMIA.17,18 Figure 3 lists specific country membership in standards organizations for LIC, LMIC, and UMIC. Figure 4 groups country membership according to geographic region. Note that while IMIA is not an SDO, it is included in these charts because its greater penetration among low-income countries and among target geographic regions makes it a natural forum for health informatics activities in these countries.

The majority of standards are available for purchase, through licensing or through SDO membership. HL7 recently announced that it intended to make a majority of its standards available for free19, but since most SDO revenue comes from standards sales or membership, new business models may be needed before other SDOs will follow suit.

Interoperability

Interoperability broadly refers to the ability for entities within a system to interact, while standards define the entities and their interactions. Defining interoperability is important because it “shapes the scope of the problem and thus the response in terms of policy, resources, and priorities.”20 The definition adopted by HIMSS provides a useful starting point:

“Interoperability means the ability of health information systems to work together within and across organizational boundaries in order to advance the effective delivery of healthcare for individuals and communities.”21

In 2007, HL7 surveyed leading health informatics institutions globally to further define interoperability and identified three broad categories: technical, semantic, and process.22 To these three, Palfrey and Gasser propose a fourth layer of “institutional interoperability” to capture interactions between societal systems.23 For the purposes of this report, we refer to the following four interoperability layers:

1) **Technical Interoperability** – the physical conveyance of a payload or the transmission and reception of information at the level of hardware

2) **Semantic Interoperability** – the ability of information shared by systems to be understood
3) **Process Interoperability** – the communication of information in a time-, event-, and sequence-oriented manner to optimize and coordinate the work processes of a care team

4) **Institutional Interoperability** – the structured interaction of societal systems, involving organizational models, policy and governance, trade agreements, and systems of law

While the eHealth community has prioritized technical and semantic interoperability for some time, process interoperability was not formally adopted as a standardized approach by SDOs until 2007. Institutional interoperability is often beyond the scope of purely eHealth-oriented interoperability discussions, but is included here, as many gaps and recommendations pertain to agreements between the global, regional, sub-regional, national, or sub-national institutions that shape the eHealth ecosystem.

The Integrating the Healthcare Enterprises initiative (IHE) proposes that Security be included as a distinct interoperability layer (separate from institutional or legal), given its critical importance in protecting and enabling interactions. While security is not broken out into a separate layer in this report, the need for an integrated security framework for mHealth, telehealth, and eHealth to enable interoperability is echoed throughout the literature and in stakeholder interviews.

**eHealth Architecture**

mHealth at scale must rely heavily on other health information systems, such as electronic health record systems, electronic facility registries or health management information systems. As such, mHealth must be considered within the context of the broader eHealth architecture. The approach taken in this report is that mHealth is the subset of eHealth that includes mobile phones, other mobile devices, and services delivered via telecommunications networks, the software, platforms, devices, and infrastructure that are used by the mobile device to support health system activities, and all supporting eHealth systems and standards. eHealth is defined more broadly as the application of information and communications technologies to health service delivery and supporting health system functions.

ISO TC 215 published the Technical Report (TR) 14639 “Part 1: Capacity-based eHealth architecture roadmap: Overview of national eHealth initiatives” in August 2012. This important document has been widely used by the Public Health Task Force of ISO as it presents case studies of the national eHealth architectures implemented by Australia, Brazil, Canada, India, and Kenya, and describes a national eHealth architecture framework, nicknamed the “Parthenon” (see Figure 5). The Parthenon was derived from an original graphic developed by Dr. Richard Gakuba, National eHealth Coordinator for Rwanda, as part of the Rwanda National eHealth Architecture initiative. The document recognizes that international standards typically describe systems at a high level of maturity and advocates for the development of a roadmap to guide countries in moving from a low to medium to high level of maturity. ISO TC 215 is expected to publish “Part 2: Architectural components and maturity model” in early 2013. Concurrent national eHealth architecture efforts in Brazil, Rwanda, and South Africa utilized similar frameworks as the one presented here. It is a key recommendation of this report that the framework presented in ISO TR 14639-1 be adopted by the international community to structure dialogue, investments, and development of mobile and electronic health components. Note that as an ISO standard, this report is available for purchase, although ISO TC 215 has submitted a request to the ISO central secretariat to make this standard available free of charge.
**HEALTH PROCESS DOMAIN COMPONENTS**

**Patient Path – supporting continuity of care**

- Community-based services
- Primary care services
- Hospital/ institutional services
- Public health & disease surveillance
- Emergency response
- Diagnostic services
- Pharmacy services
- Healthcare supply chain
- Human resources in health
- Health finance and insurance
- Vital records collection & management
- Environmental monitoring
- Knowledge management & eLearning

**HS planning, monitoring & evaluation**

**FOUNDATION – eHEALTH INFRASTRUCTURE**

- EHR & health information repositories
- Identification registries & directories
- Clinical terminology & classifications
- Data interchange interoperability & accessibility
- Consent/ access control & workflow management
- Privacy, security & safety regime
- Census, population information & data warehouse

**FOUNDATION – ICT INFRASTRUCTURE**

- Local access to ICT equipment and facilities
- Electronic communications infrastructure
- ICT processing and storage services
- ICT professional & technical support
- Standards, methods, guidelines, frameworks

**GOVERNANCE & NATIONAL OWNERSHIP**

- Executive sponsorship
- National leadership of eHealth program
- eHealth standards adoption & implementation
- Development of eHealth capability & capacity
- eHealth financing & performance management
- eHealth planning & architecture maintenance

*Figure 5. National eHealth Architecture from ISO Technical Report 14639-1 – Capacity-based eHealth architecture roadmap – Part 1: Overview of national eHealth initiatives.*
Literature Review

Overview

The following literature review was performed to assess how mHealth standards and interoperability have been addressed in academic and gray literature, and to identify the gaps.

The academic community plays an important role in evaluating the benefits of employing mobile and electronic health systems to facilitate the delivery of health services, and there is a growing body of evidence to support their use.26 Yet, evaluations of the comparative value of adopting specific health informatics standards in support of mHealth programs were uncommon. This is not unexpected, however, as there are very few scaled-up, interoperable implementations to be evaluated at the time of publication.

While there has been an upsurge of scientific articles in the past decade on mHealth standards and interoperability, the majority were focused on high-income settings and narrowly on telemedicine or home telehealth. Peer-reviewed articles that looked at LMIC settings were generally descriptive in nature, often advocating for global alignment around eHealth standards, interoperability, or capacity issues.62 However, a scan of gray literature uncovered numerous LMIC-focused documents describing international, national, or sub-national efforts to support local actors in LMICs to tackle standards and interoperability challenges.

The review indicates that the integration of mHealth into the eHealth ecosystem is an inadequately defined problem space. The issue is exacerbated by the realities of working in developing health systems, which often includes limited resources, poor infrastructure, and the absence of an existing eHealth infrastructure with which to integrate.

Analysis of Literature and Systematic Reviews

Three literature reviews addressed specific sub-topics within mHealth standards and interoperability: (1) health informatics for development; (2) telehealth security standards; and (3) home telehealth information system architecture.

Marcelo, 2012 – Health Informatics for Development

A literature review in 2012 by A. Marcelo, A. Adejumo, and D. Luna (hereafter Marcelo) examined efforts to build health informatics capacity within low- and middle-income countries and proposed a roadmap to “fast track” uptake of health informatics for development.27 The review identified 77 articles that fit the search criteria of “health informatics” and “development”, of which only 9 focused on African or Asian countries.
The authors described the state of health informatics capacity in three regions:

- **Latin America** is generally led by the private sector, rather than ministries of health, leading to poor coordination between health informatics stakeholders and proliferation of proprietary systems. Argentina, Brazil, and Uruguay have medical informatics training initiatives, and Ministries of Health from Brazil and Chile have formally engaged in standards development and adoption, but the rest of the region has progressed more slowly.

- **Africa** has experienced an increase in health informatics activity in recent years due to increased interest in ICT projects, but implementations are fragmented, small, and generally led by the donor community rather than local stakeholders. Poor infrastructure, including unreliable electric power, is a primary obstacle. Africa is among the fastest growing mobile phone markets in the world.

- Despite presence of local expertise, governments in **Asia** have been slow to adopt recommended health informatics standards. Geography, diversity of languages, and limited investments into health informatics all contribute to slow adoption and the lack of regional efforts.

**Marcelo** described the cyclical nature of how poor infrastructure limits the growth of local capacity, which in turn limits infrastructure development. Use of proprietary software systems, lack of national privacy frameworks, and lack of standards adoption were identified as key obstacles to the development of local health informatics ecosystems.

**Marcelo** proposes a three-pronged strategy to fast track the development of local health informatics capacity: establishing partnerships, adopting standards, and leveraging mobile health. **Partnering** with established global health informatics institutions (e.g., a collaboration agreement between a local and foreign university) will accelerate the rate of knowledge transfer to low- and middle-income countries. Pairing training with implementations will ensure that informatics practices are contextualized and incentivize involvement for both parties. One notable example is the Pan–Asian Collaboration for Evidence-based eHealth Adoption and Application (PANAceA) formed by the International Development Research Centre of Canada (IDRC) and Aga Khan University. PANAceA has funded eight projects since February 2007 to generate evidence for the use of eHealth within the Asian context.

The second recommendation made by **Marcelo** is to adopt standards for interoperability during the formative period of the ICT infrastructure and health informatics ecosystem. Mature standards did not exist when high-income economies began developing eHealth systems, which has contributed to fragmented systems and poor interoperability. LMICs have the opportunity to avoid this by proactively defining their unique functional requirements and developing enterprise architectures to guide ecosystem development. The exchange of indicator definitions and data for aggregating reporting, possibly using the SDMX–HD standard developed by the WHO, and standardized geographic codes are emerging standards categories that have the potential to impact developing world health systems.

The final recommendation made by **Marcelo** is to leverage mobile phones for health. The pervasiveness and well-understood interfaces of mobile phones provides new opportunities for decentralized data collection and reporting.
Garg and Brewer, 2012 – Telemedicine Security Standards

Garg and Brewer, 2012\textsuperscript{19}, in February 2011, performed a systematic review of 58 articles raising serious concerns around security and privacy standards utilization within telemedicine. Articles represented 11 high-income and 2 middle-income countries. Despite the availability of mature privacy standards, including the Health Information Portability and Accountability Act (HIPAA) in the United States and HL7 that is widely used in Europe, not a single article specifically tested security. 61\% of articles proposed solutions for specific security issues, but few exploited a comprehensive security framework, which must address privacy, data integrity, authentication, authorization, and reliability. A limited number of articles provided details of security protocols implemented, and several that did so used protocols that were outdated or that had known flaws.

Garg and Brewer note that telemedicine presents some security requirements that are not common to other forms of health information technology, in particular the reliability, availability, and usability of systems. These issues must be addressed before telemedicine can be consistently used in lieu of in-person clinical encounters. Telemedicine researchers are encouraged to collaborate closely with security experts to address numerous gaps in telemedicine security.

Ludwig – Home Telehealth Information Systems Architecture

Standards organizations have not yet endorsed a reference architecture for home telehealth information systems. Ludwig \textit{et al} (hereafter “Ludwig”)\textsuperscript{29} reviewed 22 articles to develop a nomenclature for the information and communications technologies that support home telehealth services as a preliminary step in development of a reference architecture. Architectures to support telehealth information systems differ from traditional architectures in two important ways:

- Home telehealth services are \textit{sensor-enhanced}. This means that devices in the patient’s home or those worn by the patient are automatically generating data and integrating that data into a health information system.
- Home telehealth services are \textit{trans-institutional}, which means that two or more distinct legal entities are involved in managing a patient’s care, raising questions about ownership, security, duplicative data stores, and assignment of responsibility.

Ludwig presents a nomenclature broken into six categories: users, services, operating organizations, information flow, geographical reach, and architectural paradigm. While this nomenclature targeted home-telehealth within high-income countries, the nomenclature may be expanded to incorporate other aspects of mHealth or provide a template for developing a similar nomenclature within specific mHealth subsets.
Supplementary Articles Review

In total, 2,154 published articles from peer-reviewed sources matched the above search criteria, although the majority of articles were duplicative or fell within the exclusion criteria. After screening, 46 published articles met the search criteria.

It is important to note that there were a large number of articles covering the topics of eHealth standards and interoperability without explicit mention of mobile health or a variation of the term. A search on PubMed for eHealth related terms with a standards or interoperability term resulted in well over 5,000 articles prior to screening. Many standards previously used exclusively in the context of eHealth are also applicable within mHealth (e.g., standardized vocabularies, security protocols, etc.), while others require adaptation to support different form factors or use cases. Given the overlap between eHealth and mHealth, it is probable that many articles that are pertinent to the topic of mHealth standards and interoperability were not identified due to this difference in classification.

The goal of the review was to explore how mHealth standards and interoperability were being addressed in scientific literature. The breakdown below provides a snapshot:

- 3 articles were literature reviews looking at (1) health informatics for development (2) telemedicine security standards and (3) home telehealth information system architecture. No crosscutting literature reviews were found that examined mHealth standards and interoperability more broadly.
- 24% of published articles targeted low- and/or middle-income countries or were global in scope, 63% targeted high-income countries, and 15% addressed technical standards with no specific country target. LMICs referred to specifically included Philippines, Uganda, and Uruguay (note that articles may fall in more than one category). For an LMIC example, Margolis et al present a countrywide EHR implementation with prominent use of health informatics standards, including terminology services and telemedicine applications.\(^30\)
- 9% proposed a new standard or assessed the relevancy of a standard based on a specific use case.\(^31\)
- 15% proposed or described an implementation of one or more standards for a specific use case.\(^32,33,34,35\) For example, Constantinescu et al demonstrated the use of standards-based protocols to deliver multimedia-rich content to mobile handsets from EHR, diagnostic imaging repositories, and other systems.\(^36\)
- 30% of articles focused on mHealth security or privacy. DD Luxton et al also identifies the lack of security standards as a key obstacle to supporting “ubiquitous use of mobile health technologies.”\(^37\) P. Pharow et al advocated for development of a standards-based interoperable security infrastructure.\(^38\) Koster et al “introduced” end-to-end security requirements and [presented] a design for consent management in the context of the Continua Health Alliance architecture.\(^39\) Adoption of standards supporting identity management within eHealth and telemedicine are far behind the maturity of the standards.\(^40\)
- 2 articles specifically recommended that the health informatics community revise the definition of interoperability to better address an overarching security framework in the context of interconnected systems.\(^24,41\)
• 9 articles (20%) made explicit mention of HL7 standards, 3 (7%) of IHE profiles, 4 (9%) of DICOM, and 6 (13%) of ISO/IEEE 11073 family of standards for medical device interoperability.

• 3 articles proposed chronic disease, in particular diabetes, as the use case for interoperability within a heterogeneous environment of patient or home-based health devices.

• 6 articles focused on development of home telehealth networks capable of integrating information from heterogeneous sources, including personal health devices and sensors.

• 2 articles did the challenging work of proposing how a group of related standards can be used in conjunction to achieve interoperability within a heterogeneous environment of health systems and devices. Mense et al proposed development of a framework for leveraging international standards ISO/IEEE 11073, HL7, IHE, CEN 13606 to integrate data from a mixed environment of proprietary devices.

• Several examples of standards-based software components and platforms specifically address the issue of intelligently integrating information from multiple personal health devices, including mobile phones. Middleware components were the most common integration architecture proposed. While these tools employ standards, many of the interfacing components and related processes remain proprietary. Special care is needed to handle variance in the accuracy, bias, and availability of data from these sources.

• Blobel presented an architectural framework to support a lifelong patient record that is scalable, flexible, portable, secure, and integrates data from heterogeneous sources. The architecture must follow the ISO reference model for open distributing processing (ISO RM-ODP). Examples include Australian GEHR, openEHR initiative, CEN ENV 13606 “Electronic Health Record communication”, and HL7 version 3 (specifically HL7 v3 Reference Implementation Model, HL7 Development Framework, and HL7 Clinical Document Architecture).

• While no articles evaluated the patient or health system impact of the use of standards or systems integration, Michael et al described the design of an impact evaluation of mHealth within the existing eHealth architecture already in use by the Millennium Villages Project.

As expected, the number of relevant articles increased in recent years, with 81% of the articles being published after 2007 (Figure 3).
The Telemedicine Alliance, a consortium comprised of the ESA, WHO, and ITU formed in 2002, identified interoperability as the "main showstopper" in eHealth implementation. Indeed, the preponderance of articles reviewed stated the need for mHealth standards and interoperability in order for mHealth projects to lead to improved health system performance at scale. Specific concerns included:

- Lack of standards compliance or awareness
- Lack of technical compatibility or interoperability between systems
- Lack of mechanisms for collaborative investments into eHealth
- Inadequate local health informatics expertise
- Widespread use of proprietary systems and databases
- Reference architectures and ontologies lack new mHealth use cases
- Increase in the volume and quality of data as devices used by frontline workers (FLWs) and patients are integrated into health information systems

M. Ackerman et al also identified interoperability of data among telehealth systems and integration with other electronic systems, such as EHR, as key obstacles to effective use and analysis of telehealth, and a potential source of dissatisfaction among patients. The article makes several recommendations to the US National Institutes of Health that may be translatable to governmental institutions in resource-constrained settings:

- Conduct multi-stakeholder meetings with focused deliverables and provide funding opportunities for convening
- Fund innovative research to develop, assess, and support telehealth-related tools and technologies, with a focus on translational platforms
- Fund implementation research to improve delivery, use, and adoption of telehealth technologies
- Create a central information resource compiling free telehealth software and resources
Key Informant Interviews

Key informants were interviewed to inform the gaps analysis and to gauge how each stakeholder group uniquely perceives and engages with mHealth standards and interoperability.

Results

1. Perceptions of the past and future of interoperability efforts

Interoperability efforts are not new and opinions on the effectiveness of past efforts were mixed. A common sentiment was that interoperability has turned out to be far more challenging than expected. Some key informants believed that in the past, there was not a strong enough central force to achieve the necessary coordination across a variety of stakeholders. Several cited that the lack of a common definition for interoperability and the lack of common measures for success for interoperability contributed to an overall perception that interoperability efforts were ineffectual. Some informants pointed to the fact that there are few examples of major eHealth and mHealth projects that are interoperable. There was concern that this may lead to lower participation and reduced confidence in the eventual success of future high-level efforts.

There was an alternative view that prior interoperability efforts were successful, just not in the way that was expected. As one informant said,

“When the conversation first started, it was trying to explain what interoperability was and what a standard is and why these are important...but now it is almost universally accepted that interoperability is needed. You have governments writing this into their strategies.”

The sentiment was that the conversations of the last decade had, in a way, set the stage for today’s major interoperability efforts, such as those seen in the Rwanda Health Enterprise Architecture initiative and the MoTech Suite.

Interoperability was on the agenda of stakeholders of all types, from governments and standards developers, to vendors and donors. The most common need for interoperability was that tools could share critical health information. For most stakeholders, the drivers behind this were: the difficulty interfacing systems of different types, from different vendors, or collected from different health worker types; aggregating individual patient data for district- or national-level reporting; and comparing data from several similar projects by different implementers funded by different donors. Donors and implementers were also interested in the ability to repurpose software components developed in one setting in another context.
Among vendors, the idea that systems should interoperate has resistance from some entities who have legitimate competitive or cost-effectiveness interests. In some cases, market pressures actually reward non-standards-based efforts for differentiating themselves. Innovation grants may inadvertently encourage new developments or incompatible technologies, and they rarely carry the budgets necessary to support standards compliance. In a competitive market, vendors can achieve lock-in by intentionally not adopting standards.

As one key informant noted, “interoperability is a reflection of social and institutional relationships,” yet many initiatives in the space have focused solely on the technological aspects. Another informant commented, “There is no one who has any degree of interoperability accidentally.”

Lack of clarity of the necessary steps and the costs of achieving interoperability were commons themes.

2. Perceptions of health informatics standards

“Standards are a bit like doing plumbing – technically challenging, critical to get right, no one wants to do it.”

While all key informants agreed that standards play a valuable normative role, there was a wide range of opinions of the relevance and levels of knowledge of health informatics standards. An issue cited by many informants was the perception of the lack of relevance of standards to pilots and to LMIC projects.

“There is this idea that the use cases are so different that existing standards are not relevant, at least not without major adaptation.”

The fact that LMIC countries have little representation among SDOs helped to reinforce the idea of “developing country exceptionalism.” Other informants suggested that LMIC and HIC environments are more similar than they are different.

“It’s suggesting that these environments are so fundamentally different that there’s never going to be convergence. If you look at other domains, financial markets, how they build and run airports and shipping…that doesn’t hold at all.”

Most informants agreed that increased engagement of LMICs among SDOs is needed, but that does not necessarily equate to developing new standards for LMIC-specific approaches to problems that have already been solved for HICs. An interviewee stated, “LMICs no longer start with a blank sheet of paper; they start instead with guidance.”

Informants proposed two potential solutions to this issue: one, the development of implementation guides to assist implementers in employing specific international standards; and, two, creating a compendium of standards and their use cases to help users understand which standards are relevant to meeting particular requirements. Capacity building and training on standards was recommended for ministries of health, implementers, and donors, in particular, to encourage prioritization of standards-based approaches, to increase awareness of what standards exist, and to improve stakeholders’ ability to effectively adopt.
Limited access by LMICs to published health informatics standards was a common theme, in part because the majority of standards are only available for purchase and, more importantly, the cost of adoption is high. Interestingly, most informants were not aware of the price of standards (which can be as little as 100 USD, although many are much more expensive). Many key informants called for “open standards” and stated the need for global leaders to step up and advocate for standards to be made available for free. As one informant stated, “Openness is not historically a part of the international standards development process,” and only recently have major SDOs taken this issue on. One informant recommended an interesting alternative that may bypass the issue — development of open-source software components that are based on closed international standards, but made available as public good.

Key informants recognized the difficulty of balancing between pragmatism and standards-compliance. Ideas that are common in mHealth, such as pilots, prototyping, minimum viable products, and quick iteration of software, were all seen as being in competition with standards-based approaches, which require careful architecting and design. Others suggested that going through the process of developing or implementing a homegrown solution helps people understand the need and value of standards-based solutions. One technology and standards developer acknowledged that it often does not make sense to adopt complex representations described by international standards first. While there are “lightweight” versions of certain standards that exist just for this purpose (e.g., Green CDA), informants agreed that more lightweight versions are essential if standards are to become widespread for mHealth and among LMICs.

One key informant suggested that the split between pragmatism and standards-compliance was a false dichotomy, and that you should always “optimize for pragmatism.” In other words, the realities of a specific business need and context must always be accommodated, while still meeting an acceptable level of compliance with international standards. Key informants noted the lack of appropriate forums to discuss balance points with the broader community.

While there was the expectation that some new standards are needed and that some existing standards will need to be adapted to support mHealth within LMICs, key informants stated that an eHealth reference architecture that adequately addressed mobile clients is the critical first step since it serves as a framework for future standards work. Stakeholders from SDOs made it clear that until a mHealth reference architecture is developed, it will not be clear what the gaps are in health informatics standards in order to enable mHealth to interoperate with other health information systems. From the HL7 standpoint, this means identifying the linkages between the mHealth architecture and the existing Electronic Health Record, Personal Health Record, and Public Health profiles.

3. mHealth relative to eHealth

There was unanimous agreement that the global community should not address mHealth independently from eHealth. Two notable quotes:

- “There is no mHealth without eHealth.”
- “We do not make a distinction between m and e in terms of health data standards.”
Yet, there was lack of clarity about the actual standards and interoperability issues that are particular to mHealth. In other words, steps should be taken to determine how to approach the broader eHealth ecosystem through the lens of mHealth. This was regarded as a “critical” or “central” issue by several informants.

Informants from SDOs concurred that they had yet to agree on what new standards development or adaptation was needed to accommodate requirements for mHealth usage worldwide, but they also did not necessarily see this as a unique problem. As one SDO informant stated, each actor throughout the health system will have their own definition of what mHealth is, and it may not be the same from country to country. Elucidating ambiguities and developing consensus on the definition of mHealth from each particular stakeholder’s point of view is part of the process of developing a mHealth reference architecture.

4. The pivotal role of donors

Donors were seen as being key drivers of eHealth interoperability in the future, but also as having contributed to the existing fragmented eHealth environment. One key informant put it very succinctly:

“If there is always funding for a silo, then a silo will always be built.”

Some interviewees placed much of the burden for a shift towards eHealth interoperability on donors because of their ability to influence market dynamics and incentives in favor of platforms, vendors, governments, and implementers that agree to employ standards and meet specified interoperability requirements. One mechanism proposed was to include language in requests for proposals (RFPs) requiring demonstrated compliance with interoperability requirements or adequate justification for non-compliance.

While this sounds promising, others were skeptical, noting that interoperability is only funded when it is needed to support a specific health problem, meaning that interoperability is not often funded directly. Two interviewees pointed to the fact that no donor has funded the development of a robust interface between mobile phones used by community health workers and commonly used electronic health record systems, despite the fact that it was defined as an important obstacle several years ago. Some informants noted that donors do not often have sufficient expertise in standards and interoperability to accept much risk in their investments into the space. As one donor said:

“We’re not very sophisticated on [interoperability]. We have no in-house competency in interoperability and do not have a clear position on it...We expect those that we fund to be able to make decisions about interoperability.”

The lack of coordination and alignment between donors was a common topic. An informant brought up the common example of two projects addressing the same health priority within the same ministry of health being entirely incompatible because separate donors funded them. A donor acknowledged that there are no accepted modalities to jointly fund interoperability efforts at the ministry level across multiple independent donor projects. Many informants recognized the need for a global forum for donors to achieve alignment amongst themselves and with governments, SDOs, and implementers.
Three informants proposed independent certification of mHealth and eHealth software as a possible mechanism that donors could use to ensure their investments met objective measures for standards and interoperability compliance. In addition, funding of “camps” or “product suites” that can be marketed as integrated packages was suggested as a pragmatic way forward. This would be similar to the lines of products from Apple or Microsoft, which have very little interoperability with each other, but are highly integrated internally.

5. Role of other stakeholders in health informatics standards uptake

Each stakeholder has different priorities, incentives, and capacity with respect to standards and interoperability.

Governments were considered to be the primary target for standards adoption efforts, as they are the primary organizational beneficiaries of interoperable systems and they ultimately have the potential to serve as gatekeepers for eHealth deployments within their health systems. Interviewees acknowledged that there are many obstacles to LMICs effectively adopting standards and interoperability best practices. Specific obstacles included: a general dearth of experience and expertise with standards; lack of prioritization and understanding of the need for standards; limited resources for standards activities; under-empowered national standards bodies (if they exist at all); and little participation in the international standards process. One government representative said:

“I can’t overemphasize the need for governance, for eHealth strategy, standards, and implementation monitoring [compliance]. That has been our greatest thorn here...This is where ministers [not abstract ministries] must take leadership [or must actively delegate] or else any eHealth governance activity will fizzle out...”

Several interviewees lauded the trend towards developing national eHealth strategies among LMICs with appropriate mention of standards and interoperability, but noted that many countries are struggling to operationalize these strategies due to the same issues listed above. While international efforts to increase uptake of standards among LMICs will be helpful, several indicated that there is no replacement for national forums to make standards relevant to country efforts. Given the influence donors have on national health systems, donors must align their activities and RFPs with countries’ actual eHealth priorities.

The technology developers and vendors tended to be very familiar with important information technology standards, such as JSON, XML, or REST, but less familiar with health informatics standards, even in cases where those standards are built off the same technology (i.e., JSON, XML, and REST). The technology developers tend to be focused on a single platform, and, as one developer said, “standards are only considered relevant when interfacing with another information system,” which is typically the responsibility of the implementer or government, not the technology developer.

Several informants noted that many of the interoperability issues are technologically simple, but that there is no one directly funding it. Referring to designing an interface between mobiles and electronic medical records, one developer commented that if a donor required it in order to apply for a RFP, or if it were funded directly, “it could be ready in a month.”
Even if funding were available, the market incentives may encourage development of non-interoperable systems. One informant noted:

“As long as this is true that if I have more money and can go faster it will become a de facto standard, then those seeking market share or product stacks will have a disincentive to adopt.”

The competitive advantage to building proprietary systems may never go away, but governments, donors, and other payers have the ability to shift market dynamics in such a way that compliance is rewarded, for example, by facilitating access to a larger customer base. In some cases, this actually lowers risk for vendors—and implementers—by providing assurance of long-term market compatibility.

**Implementers** most frequently cited real world mHealth interoperability struggles, as they often have the responsibility for making multiple systems work on the ground. Implementers have been quicker to adopt innovative mhealth solutions than governments, which means they have felt the growing pains of a nascent field. Implementers have also been among the first to adopt standards and to work towards interoperable systems. Implementers cited issues of insufficient capacity, lack of awareness of relevant standards for interoperation, limited mechanisms to align technology approaches with government or other local implementers, and inability to divert limited grant funding towards development of generic software interfaces.

**Academics and researchers** noted that the lack of common terminology and interoperable systems was a key obstacle in comparing results across projects, thus limiting their ability to perform evaluations. The paucity of existing interoperable deployments has prevented the evaluation of the comparative effectiveness of adopting health informatics standards.

**Multinational organizations** were most concerned with aggregation of health information from heterogeneous sources and challenges of coordinating a large community of diverse stakeholders.
Discussion of Gaps and Opportunities

The gaps and opportunities with respect to the use of standards to achieve interoperable mHealth at scale can be categorized into five buckets: (1) Building capacity and increasing standards access; (2) Filling in standards gaps for LMICs; (3) LMIC engagement and mediation; (4) Promoting coordination and alignment; and (5) Strengthening mHealth and eHealth governance.

1. Building capacity and increasing standards access

Building up of health informatics capacity and ensuring access to international standards are critical components of an overarching strategy for the increased uptake of health informatics standards and interoperability best practices. Each stakeholder involved in mHealth has capacity needs, but the shortage is felt more acutely by LMIC governments and local implementers. Capacity can only be built through a mix of short- and long-term efforts.

**Establish formal health informatics training and education programs** – The global eHealth community can support institutionalization of health informatics programs at universities and technical colleges within LMICs, and establish scholarships to degree-granting programs or certificate courses at foreign institutions. Formal training programs can be supplemented by online eHealth technical training courses around interoperability, standards, tools, implementation guides, case studies, etc. Education is a long-term approach. In the short-term, a mechanism for providing access to international health informatics expertise could ameliorate the workforce shortage.

**Inform local decision and policy makers of the implications of adopting (or not adopting) standards and interoperability best practices** – While this constituency does not need technical standards training, better information is needed about the costs, best practices, and governance of mHealth and eHealth infrastructure. LMIC governments will need to prioritize national standardization and harmonization activities, in part by empowering and funding national standards bodies. Decision makers also need increased awareness of which standards can be employed for particular scenarios and guidance on how to select between multiple options and adapt for local needs. Implementation guides will be a key component of this. Coupled with in-country workshops and regional events organized in collaboration with the WHO, SDOs, and other key stakeholders. The WHO is pursuing a global resolution on health information sharing and standards harmonization that will help guide policy makers in this process.

**Improve donors and implementers’ ability to strategically invest in an interoperable health system** – Similar to local policy makers, donors, implementers and other stakeholders are also in need of additional capacity in order to make more strategic investment decisions and to select appropriate standards and platforms. Many of the same strategies can be used with this group.
Empower local professionals and professional organizations – Professional organizations, such as IMIA, play an important role in building and retaining local expertise. Road shows hosted in LMICs organized in conjunction with a consortium of SDOs and professional organizations may be an important avenue for linking to local industry.

Address barriers to standards access – There is inadequate access among all stakeholders to information with respect to selection, adoption, deployment, and maintenance of standards. Part of this issue can be solved by encouraging governments to provide actors within each of their countries with national requirements, norms, and guidance for design and implementation of eHealth systems. Donors should also provide similar guidance to their grantees. But before this guidance can be provided, certain information gaps need to be filled.

A global, curated compendium of health informatics standards and implementation guides relevant to LMICs would streamline access to additional information about standards. Additional implementation guides tailored to LMIC use cases would provide clarity around the use of standards within new contexts.

Even with better information, certain standards are out of reach to certain stakeholders due to their cost. The global community should advocate for other SDOs to follow HL7’s lead in open sourcing their intellectual property and collaboratively explore the development of business models to support ongoing SDO activities. Donors can also fund the development of open source software components of specific standards and the tooling to facilitate their deployment. These components could be contributed to the mHealth commons at no expense.

2. Filling in standards gaps for adoption among LMICs

Most existing health informatics standards can be repurposed without alteration in LMIC contexts, although for some adaptation will be necessary. In addition, a limited number of new standards may be needed to support unique LMIC requirements. There are three primary drivers of adaptation and possible development of standards:

1) Limited bandwidth and infrastructure leading to power fluctuations and intermittent cellular connectivity may require lighter-weight and more reliable transmission protocols.

2) The use of insecure and technically limited messaging interfaces, especially SMS and USSD, to communicate with patients, health workers, or information systems may require new methods for securing, verifying, and efficiently transmitting data and protecting patient privacy.

3) The shift to non-clinical and community-based workers has revealed gaps in standardized terminology to represent community-based care, especially in rural settings, and presents unique challenges in terms of connecting systems to enable continuity of care.
As noted in the interviews, standards adaptation and development must be guided by a mHealth reference architecture developed by a consensus process among SDOs. The reference architecture will describe the functional components of a mHealth ecosystem that links into existing functional models, such as the Electronic Health Record profile, and will comprehensively represent relevant use cases and scenarios. This work is already underway among newly created mHealth Working Groups within several SDOs, but there are areas where contributions from the global community are needed. Descriptions of existing national eHealth architectures, compilation of existing mHealth projects and their business requirements and use cases, and a more comprehensive understanding of the use of existing standards and their gaps will help inform the development of an overarching mHealth functional model. The following snapshot represents several use cases unique to projects in LMICs that merit integration into such an architecture:

- Use of short messaging service (SMS) for automated alerts, querying of patient information, patient registration, and other basic health services
- Use of SMS and Interactive Voice Response (IVR) to contact clients regarding appointment and medication reminders and lab results
- Use of SMS printers or other cellular connected devices to facilitate ancillary health system services
- Use of voice calls and SMS to send information for supply chain and inventory management
- Phone-based decision support protocol that guides an auxiliary nurse midwife through provision of antenatal care in a pregnant woman’s home

ISO TR 14639 Capacity-based eHealth Roadmap provides an important reference point for national health information systems by documenting the national health information system architectures employed in five different countries. It also contains a list of standards most relevant to such systems. Part 2 of this standard, expected in early 2013, will describe a maturity model for use of standards at low, middle, and high levels of maturity.

The standards development community is engaged in the ongoing evolution and development of standards for vocabulary, content exchange, transport, privacy, and security with respect to eHealth. It will also be necessary to add new terminology for requirements that are unique to the developing world, in particular to accommodate the significant shift towards nonclinical and community-based workers. For example, there is currently no standardized way to capture the number of nights that the members of a family slept under a treated bed net in a given period of time.

While not a technical specification, the standards community also develops implementation guides, which describe the application of one or more standards to a use case and set of business requirements that is in agreement the relevant reference architectures. An example of this is the IHE Pharmacy Profile. These guides are a critical part of ensuring that implementers employ the right standards and that governments ensure they are meeting service delivery requirements. The global health community should fund and coordinate with SDOs the development of implementation guides applicable to the implementation and integration of mHealth among LMICs.
3. LMIC engagement and mediation

The end goal with respect to the standardization gap is for LMICs to establish national health standards bodies and to become active participants in the international standards development process. Their current lack of representation among SDOs means that LMICs do not have the opportunity to contribute their requirements to standards as they are developed and that they do not have the benefit of learning from people around the world struggling with similar health system challenges. A multifaceted approach is needed to ensure that standards are applicable to LMICs and that LMICs begin the long process of establishing national bodies in order to be represented in the international standards process:

- The WHO and other multi-national institutions, donors, and implementers have significant experience working in LMICs and can collate and represent LMIC requirements to SDOs. The global community can also support the attendance of LMIC representatives to SDO meetings and technical working groups.
- Regional eHealth standardization collaboratives, established by local governments in partnership with academic institutions, private sector, SDOs, and other international organizations, can serve as a mechanism for joint regional learning, engagement with the international standards community, and can act as a valuable step on the way towards forming national standards bodies.

4. Promoting coordination and alignment

Lack of alignment and the need for regularly structured opportunities to coordinate efforts were common themes during interviews and in the literature. Some mechanisms for coordination and alignment already exist, such as the Joint Initiative Council for harmonization of standards activities among SDOs. Additional mechanisms are needed to broker the social and institutional relationships needed for interoperability among and between other stakeholder types.

In particular, there is a great need to align donor investments with eHealth standards endorsed by multi-national organizations and SDOs, as well as with the national eHealth strategies of the countries within which they are funding implementations. Donors are also in need of a mechanism to facilitate coordination of strategies and collaboration on interoperable health reporting systems, and to leverage investments into common eHealth and mHealth platforms. Empowering or establishing regional standards collaboratives, as described above, or other mechanisms are needed for LMIC stakeholders to engage with SDOs, to convey unique requirements, and to increase participation in standards development activities.

Given the influential role that donors play in shaping local eHealth deployments, they represent an important conduit through which governments and implementers can be encouraged and incentivized to adopt platforms and practices that comply with a minimum set of interoperability requirements. The WHO is currently developing Donor Alignment Principles, which are a set of high-level principles to
provide guidance for eHealth harmonization and standardization. Taking that effort one step further, donors could quickly endorse use of standards by adopting standards, interoperability and data access requirements in their grant requirements. Independent certification, e.g., mHealth Interoperability Certification, would provide an objective measure of standards compliance and can instill market confidence in a platform. Different levels of certification may be appropriate to support, for example, a system that exposes its data via standard APIs but does not comply with the full reference architecture. Donors could require or favor certification in RFPs and governments could require or favor certification in local implementations to incentivize adoption by vendors and implementers. Certification requirements would also serve as a guide for future platform development and interoperability efforts.

Certain high-profile events and awards may also encourage the technical community to adopt standards and interoperability best practices. For example, the “Best of mHealth Interop Award” could be given in conjunction with periodic “Hackers Workshops” or “Connect-athons”, such as the HIMSS IHE Connect-a-thon. These can be tied on to other major events, such as the mHealth Summit. Competitions could be held to compete for micro-grants to build interfaces for key integration points identified by a set of global health partners.

5. Strengthening mHealth and eHealth governance

Even though national eHealth strategies themselves are often strong, operationalization and governance of the policies have proven challenging. As mobile health systems move towards scale, existing guidelines and strategies will need to be revised to reflect new demands across all categories of governance. The framework in ISO TR 14639 Capacity-based eHealth Roadmap identified six categories of eHealth governance: (1) Executive sponsorship; (2) National leadership of eHealth program; (3) eHealth standards adoption and implementation; (4) Development of eHealth capability and capacity; (5) eHealth financing and performance management; and (6) eHealth planning and architecture maintenance. Regulations regarding standards adoption and interoperability of mHealth and eHealth systems will play a key role in shifting market dynamics to favor interoperability. Such regulations should be developed carefully to ensure that they do not reduce innovation, but rather shift innovation towards increasing continuity of care for patients. National interoperability efforts will be strengthened if coupled with shared facilities for compliance testing and technical assistance to adopt and adhere to standards and interoperability guidelines.

An international agreement with key stakeholders could provision the development of governance guidelines and support operationalization of eHealth Strategies across the six categories of ISO TR 14639, while building off the recommendations of the WHO-ITU National eHealth Strategy Toolkit. Workshops and networks between candidate LMIC organizations and governments to facilitate organizational interoperability and experience sharing could support operationalization and capacity building.
Conclusion

Improving adoption of health informatics standards in support of interoperable mHealth tools at scale is a complex and long-term process and there is no silver bullet solution. The following three crosscutting recommendations are intended to frame subsequent dialogue around near-term opportunities for the global community to address these challenges in a more coordinated fashion:

1. **Shift market dynamics to incentivize interoperability** – Existing market dynamics do not penalize and even reward the use of proprietary and non-interoperable systems. This is caused by many factors, including lack of funding and limited guidance for standards compliance, and, in some cases, legitimate competitive interests. Governments, through leadership and regulatory mechanisms, play the biggest role in shifting market dynamics to encourage interoperable systems. Donors have the opportunity to accelerate the market shift through strategic investments that require an appropriate level of standards compliance for grantees, through a certification process. Technology partners will bear the lion’s share of the technical burden for developing standards-based software components, but the burden can be lowered through a transparent, participatory regulatory process and incentivized through increased market access and reduced market risk. Lowering the cost of standards adoption and adaptation will reduce market disincentives. This can be done through developing implementation guides and domain profiles, funding the development of reusable open-source software implementations of standards, and providing technical assistance and resources at the national and international level for selection, adaptation, and adoption of standards.

2. **Align with national eHealth strategies** – National eHealth strategies provide a country-specific framework for the harmonization of mobile and electronic health within a country in a strategic and sustainable fashion. Alignment with the strategy provides a clear roadmap to scale and compatibility with the national information systems. Investments and implementations that do not align with the national strategy risk increased health system fragmentation, reduced local buy-in, lack of interoperability, shortened product lifespan, and, ultimately, reduced impact. A broad assessment of national eHealth strategies that builds off existing case studies and inventories performed by the WHO, ISO, and others, will be an important first step in assisting stakeholders in identifying steps needed to align with local requirements.

3. **Establish regional eHealth standardization collaboratives** – Regional eHealth Standardization Collaboratives will allow governments and local stakeholders to pool resources and expertise to make national and regional standards recommendations, to facilitate shared learning, and to serve as a focal point for capacity building activities and for interacting with international standards organizations. The collaborative would be a resource for organizations implementing mobile and electronic health tools within the region. When countries are prepared, the collaborative would also serve as a launching pad for establishing national standards bodies.
mHealth is still in its early stages and very few implementations have moved past the pilot phase. As more and more deployments are scaled up, interoperability between these systems and the broader health information system will become essential in order for health data to be made available to the right person, at the right place and time. While health informatics standards are fundamental to enabling interoperability and continuity of care, both are ultimately a reflection of human institutions and their resulting processes. Standards are an essential building block, but it is up to governments, vendors, donors, and implementers to build an enabling environment that incentivizes the use of standards-based technologies to strengthen health systems and to improve health outcomes for those most in need.
# Appendix A – Key Informant Interview List

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<tr>
<th>Name</th>
<th>Organization</th>
<th>Country</th>
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<tbody>
<tr>
<td>Alex Pavluck</td>
<td>Task Force for Global Health</td>
<td>United States</td>
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<tr>
<td>Alvin Marcelo</td>
<td>Philippine Health Insurance Corporation</td>
<td>Philippines</td>
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<td>Andrew Grant</td>
<td>Sherbrooke University</td>
<td>Canada</td>
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<td>Bart Stidham</td>
<td>Thoughtworks</td>
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<tr>
<td>Beatriz de Faria Leao</td>
<td>Consultant</td>
<td>Brazil</td>
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<tr>
<td>Bob Jolliffe</td>
<td>Consultant</td>
<td>Ireland</td>
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<tr>
<td>Catherine Chronaki</td>
<td>HL7 International</td>
<td>Greece</td>
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<td>Catherine Omaswa</td>
<td>eHealth Society of Uganda</td>
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<td>Chris Bailey</td>
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<td>Switzerland</td>
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<td>Chris Seebregts</td>
<td>Jembi Health Systems</td>
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<td>Chuck Parker</td>
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<td>Craig Frederichs</td>
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<td>Frederick Kristensen</td>
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<td>Hamish Fraser</td>
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<td>Hani Eskandar</td>
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<td>Joaquin Blaya</td>
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<td>Richard Gakuba</td>
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